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Applicant: TOHO TITANIUM CO. LTD. 13-31. Konan 2-chome
Minato-ku Tokyo 105(JP)

(2) Inventor: Okudaria, Shigenori 2-9-5, Yurigaoka Ninomiya-machi Naka-gun Kanagawa-ken(JP) Inventor: Irie, Takefumi 733, Yabata, Chigasaki-shi Kanagawa-ken(JP) Inventor: Uchida, Hiroshi 2-1-15, Murota Chigasaki-shi Kanagawa-ken(JP)

> Inventor: Fukasawa, Elichi 201, Miyuki-Helghts 633, Koyato

Samukawa-machi

Koza-gun Kanagawa-ken(JP) Inventor: Yamaguchi, Masanori 889, Matsumoto Nakai-machi

Ashigarakami-gun Kanagawa-ken(JP)

Inventor: Kobayashi, Kazuhiko

511-3-403, Chigasaki

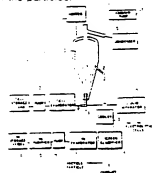
Chigasaki-shi Kanagawa-ken(JP)

Representative: Rees. David Christopher et al Kilburn & Strode 30 John Street London WC1N 2DD(GB)

Method for producing a metal from its halide.

A method for producing metal by the reduction of a metal halide by a reducing metal agent capable of continuously producing homogeneous metal of Shigh purity. In the method, particles of the same metal as the metal to be produced are charged into a reaction vessel and the metal halide and the reducing metal agent both in vapour form, are ejected upwards into the reaction vessel from its lower cortion to form a fluidized bed of the metal particles in the vessel. The reducing reaction between the two vapours takes place on the surface of the metal particles at a temperature below the melting point of the metal product and at a pressure below a vapour pressure of each of the reducing metal agent and

the metal halide at that temperature, resulting in depositing and growing the metal product on the surface of the particles.



METHOD FOR PRODUCING A METAL FROM ITS HALIDE

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The present invention relates to a method of producing metal from a metal halide and more particularly to a method for the production of a metal by the reduction of its metal halide consideration metal agent.

A conventionally known method for producing a metal by the reduction of its metal hailde comprises the so-called Kroll process, which is particularly suitable for the production of titanium. Unroducately, this process has a disadvantage in causing the metal product, for example, titanium to become firmly adhered to the side and/or bottom surfaces of the reaction vessel as a porous solid product during the production.

Another method known in the art is one in which a reaction takes place between titanium tetrachloride and magnesium on the surface of a litanium ribbon to produce titanium by depositing it on the surface. This is disclosed in Japanese Patent Publication No.3004/1958. Unfortunately, this method causes the metal product to deposit on the ribbon and also on a frame supporting the ribbon.

Accordingly, the prior art fails to provide a continuous operation for producing titanium because of the necessity to remove the deposited metal prior to carrying out subsequent steps. For this reason, the prior art methods are compelled to employ a batch process which causes discontinuity and inefficiency in the manufacturing operation. Furthermore, solid titanium or the like deposited to the inner surface of the reaction vessel is readily contaminated by the material of the reaction vessel since it is highly reactive at the elevated process temperatures, to a degree sufficient to react the reactants, resulting in deterioration in the quality and purity of the metal product.

It is therefore an object of the present invention to provide a method for the production of a metal from its metal halide which is capable of continuously producing homogeneous metal of high quanty by reduction of the metal halide.

According to the present invention, there is provided a method for producing a metal by the reduction of its metal halide using a reducing metal agent, characterized by the steps of: charging a reaction vessel with particles of the same metal as the metal product; injecting a gas upwards into the reaction vessel from its lower portion to cause the metal particles in the reaction vessel to form a fluidized bed of the metal particles; introducing the metal halide and reducing metal agent into the reaction vessel; and causing a reducing reaction of the metal halide with the reducing metal agent on the surface of the metal particles at a temperature below the melting point of the metal product and at

a pressure below a vapour pressure of each of the reducing metal agent and the metal halide at the reaction temperature.

The metal carticles serve as reaction nuclei. This men results in depositing and growing the metal product on the surface of the metal particles. The metal particles on which the metal product is deposited can be continuously tapped out from the reaction vessel.

The injected gas may comprise the metal halide and the reducing metal agent and is injected in vapour form from the lower portion of the reaction vessel to cause the metal particles to fluidize, in orger to form the fluidized bed of the metal particles. Alternatively, the reducing metal agent may be fed in liquid form to a reaction zone in the reaction vessel while the injected gas comprises the metal halide injected upwards in vapour form from the lower portion of the reaction vessel to cause the metal particles to fluidize, in order to form the fluidized bed of the metal particles. Alternatively, the injected gas may be an inert gas injected upwards from the lower portion of the reaction vessel to cause the metal particles to fluidize in order to form the fluidized bed of the metal particles.

The metal halide may be titanium chloride, zirconium chloride, silicon chloride, tantalum chloride or niobium chloride. The reducing agent may be magnesium or sodium.

When the metal halide is titanium tetrachloride and the reducing agent is magnesium, the reaction may be carried out at a temperature in the range of 1000°C to 1200°C, preferably at about 1100°C. The reaction may be carried out at a pressure in the range of 20 to 50 Torr; preferably 50 Torr.

The invention may be carried into practice in various ways and some embodiments will now be described by way of example with reference to the accompanying grawings in which the single figure is a block diagram showing a system for practicing a method according to the present invention.

The following embodiment comprises the resultation of titanium tetrachloride to metal titanium using magnesium as a reducing agent.

As snown in the Figure, titanium tetrachloride reedstock is supplied at a predetermined rate from a storage tank 1 by means of a metering pump 2 to an evaporator 3. The titanium tetrachloride is reated to vaporize and then superneated in the evaporator 3.

Molten magnesium as a reducing agent is fed from a storage vessel 4 to a magnesium ourifier 45 where it is passed through a filter, such as a titanium sponge layer so that impunties, such as

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from chromium, nickel and the like and also lively gen, nitrogen and the like remaining in the machesium may be satisfactorily removed. The contest moiten magnesium is then fed to a magnesium evaporator 7 by means of a pump 6 and heares or vaporize.

During these operations, the titanium 'erach ride evaporator 3 and the magnesium erachtation may be kept substantially at an atmospher objects sure.

The titanium tetrachloride and magnesiam accours are then supplied at a predetermined ate through pipes 8 and 9 to a reaction vesser 12 and ejected from the bottom of the reactor vesser 12 through separate injection pipes 8 and 9 ato the reaction vessel 10.

The reaction vessel 10 is also supplied with a predetermined amount of titanium particles, which are to act as reaction nuclei from a hooper 11 located above the reaction vessel 10. The titanium particles are selected from commercially available titanium powders, fine titanium particles obtained from the production of titanium sponge, titanium chips produced from cutting titanium ingot and then further powdered.

The reaction vessel 10 is kept at a temperature below the melting point of the metal to be produced, and is evacuated to a lower pressure which prevents magnesium and magnesium chloride from being condensed at the temperature. The temperature in the reaction vessel 10 is preferably kept as low as possible to facilitate the selection of a material for the reaction vessel. However, the inner surface of the reaction vessel 10 is desirably kept at a temperature as high as possible and at a pressure as low as possible for the purpose of preventing magnesium and magnesium chloride from condensing on the inner surface. Also, in order to discharge magnesium chloride as a byproduct of the reaction in the reaction vessel 10 and excess magnesium from the reaction system to recover them in liquid form in a condenser 15. the condenser 15 is preferably maintained at a higher pressure. In order to meet these contradictory requirements, the process of the present invention may be practiced under conditions where the temperature and pressure in the reaction vesser 10 are set at about 1100°C and about 50 Torr. respectively.

The titanium tetrachloride and magnesium vapours supplied to the reaction vessel 10 are ejected upwards from the lower portion thereof because the reaction vessel 10 is maintained at a lower pressure, so that the energy imparted by the vapours due to their injection causes the titanium particles charged in the reaction vessel 10 to fluidize in the reaction vessel 10.

This fluidization of the titanium particles in the

reaction resset 10 may also be carried but by spwardly ejecting an nert gas such as argon from the lower portion of the reaction lessel 10 rather than using the energy imported by the titanium retrachioride and magnesium labours, to form a fluidized bed of the titanium particles in the reaction ressel in this case the ritanium tetrachloride and magnesium vapours may be ejected into the fluidized bed of the titanium particles, and then the magnesium somewhat in excess as compared to the stoichiometric amount may be supplied to the reaction vessel 10 in order to complete the reaction of the titanium tetrachloride and magnesium.

Injection of the titanium tetrachicride and magnesium vapours into the fluidized bed of the titanium particles in the reaction vesser 10 causes the intensive exothermic reaction between the two vapours. As a result, the magnesium reduces the titanium tetrachloride on the surface of the titanium particles which form the fluidized bed and which serve as reaction nuclei, according to the following reaction:

TiCl₄(gas) + 2Mg(gas) → Ti(soiid) +2MgCl₂-(gas)

The titanium thus produced due to the reaction on the surface of the titanium particles deposited on the surface of the titanium particles, so that the titanium particles gradually increase in diameter and weight in the fluidized state in the reaction vessel 10.

The temperature of the reaction zone in the reaction vessel is kept below the meiting point of the metal product, and its pressure is kept below the vapour pressure of both magnesium and magnesium chloride at that temperature. This causes the titanium tetrachloride and magnesium to react in the vapour phase to produce the titanium depositing on the surface of the titanium particles while it is growing. At the same time, the magnesium chloride, as a by-product of the reaction, and the unreacted magnesium are discharged in a mixed gas from the reaction system without concensing in the reaction zone.

Carrying out the present invention under conditions in which the reaction zone is set at a temperature of 1100°C and a pressure of 50 Torricauses the magnesium to be maintained in the vapour state in the reaction zone, since the vapour pressures of magnesium chloride and magnesium at this temperature are 86 Torriand 736 Torricaspectively. They are therefore discharged from the reaction zone without condensing to a subsequent cooling zone in which they are liquified or solidified for recovery.

The reducing reaction develops on the surface of the titanium particles to which the reaction product deposits, and promotes successive deposition and growth of the titanium. When the diameter of

the litanium particles has increased to from living to four times their initial diameter. The san operations are drawn out from the fluidized Tell the reaction lessel 10 continuously through a 20 ding port 12 to a cooler 13, in which the last of are water-cooled to a room temperature. The last operation particles are removed via a value operation as, for example, a rocker valve and charged to screen classifier 14, which classifies the metal capticles into a fine particle group with a diameter of 0.5mm or less and a coarse particle group with a greater diameter. The coarse particle group is returned to the reaction vessel 10 via the hopper 11 for use as reaction nuclei.

The vapour mixture of magnesium chloride by-product and excess magnesium discharged from the reaction vessel 10 is directed to a condenser 15 where it is condensed to a liquid. The liquified magnesium chloride and magnesium are then collected through a "vacuum leg" into a liquid separator 16, that are separated from each other due to the difference in their densities. The separated magnesium is returned to the reaction vessel 10 for reaction and separated magnesium chloride is recovered and fed to electrolytic cells (not shown).

In the Figure, reference numeral 17 designates a vacuum pump for reducing the pressure in the apparatus.

in a reaction system constructed as described above, vapours of titanium tetrachloride and magnesium are ejected into a reaction vesser maintained at a reduced pressure, and form a fluidized bed therein, so that the reducing reaction between them takes place on the surface of the titanium particles which act as nuclei for producing titanium. Accordingly, the titanium deposits and grows on the surface of the titanium particles without depositing on the wall of the reaction vessel, which is free from any contamination by the material of the wall of the reaction vessel. Also, in the present nvention, the titanium particles which are used as reaction nuclei inherently have a larger specific surface area. This results in a significant increase in the efficiency of the reaction. Furthermore, in the present invention, the reaction zone is kept at a lower pressure. Accordingly, excess magnesium and any magnesium chloride by-product which might otherwise remain in fine internal voides of the titanium carticles are subjected to vacuum distillation while the titanium particles are fluidized in the reaction vessel, resulting in the production of metal particles of high purity which are substantially free from magnesium and magnesium chioride.

The above process has been described in connection with the production of metal titanium, however, it is to be understood that the present inven-

from is also applicable to the production or manufacture of zirconium, silicon, tantalum, problum and the like. Also in the present invention, sodium may be used as a reducing metal agent.

The present invention will now be illustrated with reference to the following non-timiting examples:

Example 1

A reaction vessel of 20cm inner diameter and lined with titanium was charged with 12kg of titanium particles have a diameter of about 0.2 to 2mm, a surface area of about 12.6 \times 10⁻⁸ to 12.6 \times 10⁻⁶ m² and a weight of about 18.8 \times 10⁻⁹ to 18.8 \times 10⁻⁶kg. The reaction vessel was subjected to external heating and kept at a temperature of 1100°C while the pressure in the reaction vessel was reduced to 50 Torr by a vacuum pump.

Titanium tetrachloride vapour superheated to 372 °C and magnesium vapour heated to 1120 °C were separately ejected from individual injection ports to the lower portion of the reaction vessel at feed rates of 7980g/hr and 2539.2g/hr, respectively, so as to fluidize the titanium particles in the reaction vessel.

Injection of the titanium tetrachloride and magnesium into the reaction vessel in this way resulted in a simultaneous reaction, which in turn resulted in the temperature in the reaction vessel tending to rise. In order to maintain the reaction vessel at a desirable temperature, the external heating was controlled to keep the temperature at 1100°C.

After the reaction had been taking place for one hour, metal particles of 2012g were drawn out from a tapping port. A residue of magnesium added in excess amount of 20% and magnesium chloride were cooled in a condenser and then separated by a liquid separator. As a result, magnesium chloride of 8465.8g was recovered in liquid form.

The metal particles thus produced were formed of titanium deposited and grown on the surface of the titanium particles serving as reaction nuclei, and were substantially free from contamination by the material of the reaction vessel wall. Also, no magnesium chloride nor any of the excess magnesium remained in the fine internal voids of the metal particles. Thus, the metal product was very cure.

Example 2

A reaction vessel of 30cm inner diameter and used with titanium was charged with 30kg of titanium particles used in the Example 1. The reac-

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tion vessel was neated by an external neater and kept at a temperature of \$100°C while the pressure in the reaction vessel was required to \$2°C... by means of an automatic pressure controller.

Fitanium tetrachloride and magnesium labours were separately ejected from individual medium corts to the lower portion of the reaction lesser at feed rates of 17 95kg/hr and 5.7kg/hr, respective is on as to fluidize the titanium particles in the reaction vessel.

Injection of the titanium tetrachlorice and magnesium into the reaction vessel in this way resulted in a simultaneous reaction, which in turn resulted in the temperature in the reaction vessel tending to rise. In order to maintain the reaction vessel at a desirable temperature, the external heating was controlled to keep the temperature at 1100°C.

After ten minutes, metal particles were started to be drawn out from a tapping port at a rate of 5.43kg/hr. After cooling, the metal particles thus produced were subjected to screening, and particles of 0.5mm or below were returned to the fluidized bed from the top of the reaction vessel at a rate of 0.9kg/hr.

A magnesium chloride by-product and an excess magnesium, the total amount of which was 19.0kg/hr, were condensed in a condenser maintained at 777 °C and obtained in liquid form through a vacuum leg.

The reaction was continued for five hours. The metal particles thus produced were formed of titanium deposited and grown on the surface of the titanium particles serving as reaction nuclei, and were substantially free from contamination by the material of the reaction vessel wall. Also, no magnesium chloride nor any of the excess magnesium remained in the fine internal voids of the metal particles. Thus, the metal product was very pure.

Example 3

Example 1 was repeated except that the titanium tetrachloride and magnesium were injected into a reaction vessel in the form of gas and liquid and at feed rates of 8kg/hr and 2.6kg/hr, respectively.

The reaction proceeded in substantially the same manner as in Example 1. After two hours, the reaction was terminated and titanium particles of about 4.0kg were recovered from the reaction vessel. The particles had a particle size of 0.3 - 2.2mm and an average particle diameter of 0.8mm.

<u>Comparative</u> Example 1

A vapour chase reaction detween Ittanium tetrachioride and magnesium was conducted under substantially the tame conditions as Example 1 except that Ittanium particles were not charged into a reaction vessel initially.

It was observed that a small amount of litanium was produced on the wall of the reaction vessel. However most of the product was discharged to a concenser in the form of lower phiorides of titanium without forming any metal titanium.

Claims

- 1. A method for producing a metal by the reduction of its metal halide using a reducing metal agent, characterized by the steps of: charging a reaction vessel with particles of the same metal as the metal product; injecting a das upwards into the reaction vessel from its lower portion to cause the metal carticles in the reaction vessel to form a fluidized bed of the metal particles; introducing the metal halide and reducing metal agent into the reaction vessel; and causing a reducing reaction of the metal halide with the reducing metal agent on the surface of the metal particles at a temperature below the melting point of the metal product and at a pressure below a vapour pressure of each of the reducing metal agent and the metal halide at the reaction temperature.
- A method as claimed in Claim 1 characterized by continuously tapping the metal product deposited on the metal particles from the reaction vessel.
- 3. A method as claimed in Claim 1 or Claim 2, characterized in that the injected gas comprises the metal halide and the reducing metal agent and is injected in vapour form from the lower portion of the reaction vessel to cause the metal particles to fluidize, in order to form the fluidized bed of the metal particles.
- 4 A method as claimed in Claim 1 or Claim 2 maracterized in that the reducing metal agent is fed in liquid form to a reaction zone in the reaction lesser while the injected gas comprises the metal natice injected upwards in vacour form from the ower portion of the reaction vesser to cause the metal particles to fluidize, in order to form the fluidized ped of the metal particles.
- 5. A method as claimed in Claim 1 or Claim 2 maracterized in that the injected gas is an inert gas injected upwards from the lower portion of the reaction vessel to cause the metal particles to maidize in order to form the fluidized bed of the metal particles.

15. A method as claimed in any preceding training transferized in that the metal halide is fitanium training, striconium chloride, silicon chicride fantalum chloride or niobium chloride.

The Almethod as claimed in any preceding 13 m characterized in that the reducing metal agent is magnesium or sodium.

3. A method as claimed in any preceding 1 am characterized in that the reaction is carried out at a temperature of 1100°C and a pressure of 20 - 50 Torr.

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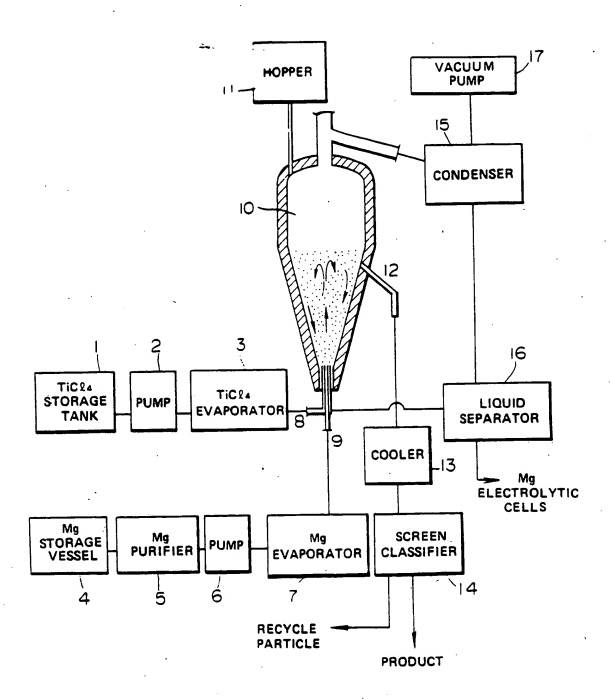
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EUROPEAN SEARCH REPORT

EP 38 30 6110

Category	Citation of document with inducation of relevant pass. 17	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CL.4)
Х . i	US-A-3 012 862 (L. 35-7-4-) et al.) * claim 1 *.	1	C 22 B 5/04 C 22 B 34/12
Α	US-A-2 827 371 (J.P. CUIN) * claims 6,9 *	1	
A	US-A-2 997 385 (C.H. WINTER, JR.) * claim 1 *	1	
A	GB-A- 736 852 (THE NATIONAL SMELTING CO.)		
A	CHEMICAL ABSTRACTS, vol. 53. 10th Jannuary - 25th February 1959, page 12145i, abstract no. 12145i, Columbus, onio, US; S. TAKEUCHI et al.: "Apparatus for continuous production of pure titanium and ziraconium", & JP - B - 58 3004 (Cat. D)	:	
	• •		TECHNICAL FIELDS SEARCHED (Int. Cl.4)
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PO FORM 1501 01 62 (P)	CATEGORY OF CITED DOCUMENTS X: particularly relevant if taken alone Y: particularly relevant if combined with another accument of the same category A: technor-gical background O: non-written disclosure P: intermediate document	earner patent do urer ine filing i D socument cited I document cited	re underlying the insocument, but published are in the application for other reasons same patent family, c	d on, of	

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